

Identification of Novel Human Kallikrein-Like Genes on Chromosome 19q13.3 - q13.4

GEORGE M. YOUSSEF^{1,2}, LIU-YING LUO^{1,2} and ELEFTHERIOS P. DIAMANDIS^{1,2}

¹Department of Pathology and Laboratory Medicine, Mount Sinai Hospital, Toronto, Ontario, M5G 1X5; ²Department of Laboratory Medicine and Pathobiology, University of Toronto, Toronto, Ontario, M5G 1X5, Canada

Abstract. The human kallikrein gene family is localized on chromosome 19q13.3-q13.4 and currently includes three members: KLK1 or pancreatic/kallikrein, KLK2 or human glandular kallikrein and KLK3 or prostate-specific antigen (PSA). The latter two genes are almost prostate-specific and they are used for diagnosis and monitoring of prostate cancer and more recently, in breast cancer applications. In this paper, we analyzed a 300Kb genomic DNA region around chromosome 19q13.3 - q13.4 in an effort to map known kallikrein or kallikrein-like genes and identify new kallikrein-like genes. Using the known kallikrein or kallikrein-like genes PSA, KLK2, zymase and normal epithelial cell-specific 1 gene (NES1) as landmarks, we have identified another six novel genes of which, five have protein homologues and gene structure similarities with other kallikreins or kallikrein-like genes. We conclude, contrary to the current belief, that the human kallikrein gene locus contains a large number of kallikrein-like genes (at least thirteen). In this paper, we present a detailed description of the human kallikrein gene locus, encompassing the already known and newly identified genes. These new genes, like the

Non-Standard Abbreviations: PSA, prostate-specific antigen; hK2, human glandular kallikrein (KLK2); NES1, normal epithelial cell-specific 1 gene; Kb, kilobase; KLK, kallikrein; KLK1, kallikrein-like; cM, centi-Morgan; EST, expressed sequence tag; RT-PCR, reverse transcription polymerase chain reaction; PAC, P1-derived artificial chromosome; BAC, bacterial artificial chromosome; TLSP, trypsin-like serine protease; HSCCTE, human stratum corneum chymotryptic enzyme; FW, forward strand.

Correspondence to: Dr. E.P. Diamandis, Mount Sinai Hospital, Department of Pathology and Laboratory Medicine, 600 University Avenue, Toronto, Ontario M5G 1X5, Canada. Fax: 416-586-8628. E-Mail: ediamandis@mshospital.on.ca

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Kallikreins and kallikrein-like proteins are a subgroup of the serine protease enzyme family and exhibit a high degree of substrate specificity [1]. The biological role of these kallikreins is the selective cleavage of specific polypeptide precursors (substrates) to release peptides with potent biological activity [2]. In mouse and rat, kallikreins are encoded by large multigene families. In the mouse genome, at least 24 genes have been identified [3]. Expression of 1 of these genes has been confirmed; the rest are presumed to be pseudogenes [4]. A similar family of 15-20 kallikreins has been found in the rat genome [5] where at least 4 of these are known to be expressed [6]. Three human kallikrein genes have been described, i.e. prostatic specific antigen (PSA or KLK3) [7], human glandular kallikrein (KLK2) [8] and tissue (pancreatic-renal) kallikrein (KLK1) [9]. The PSA gene spans 5.8 Kb of sequence which has been published [7]; the KLK2 gene has a size of 5.2 Kb and its complete structure has also been elucidated [8]. The KLK1 gene is approximately 4.5 Kb long and the exon sequences and the exon-intron junctions of this gene have been determined [9]. The mouse kallikrein genes are clustered in groups of up to 11 genes on chromosome 7 and the distance between the genes in the various clusters can be as small as 3-7 Kb [3]. All three established human kallikrein genes have been assigned to chromosome 19q13.2 - 19q13.4 and the distance between PSA and KLK2 have been estimated to be 12 Kb [9].

A major difference between mouse and human kallikreins is that two of the human kallikreins (KLK2 and KLK3) are expressed almost exclusively in the prostate while in mouse, none of the kallikreins is localized in this organ. Other candidate new members of the human kallikrein gene family include procase M [10] (also named zymase [11] or neutrosin [12]) and the normal epithelial cell-specific gene 1 (NES1) [13]. Both genes have been assigned

alignment of the known sequences of these genes with the 300 Kb contig enabled us to precisely localize all four genes and determine the direction of transcription, as shown by the arrows in Figure 1. The KLK1 gene sequence was not identified on any of these contigs and appears to be further telomeric to NES1 (since it co-localized on the same PAC as NES1). We did not attempt to characterize the genomic position of the KLK1 gene.

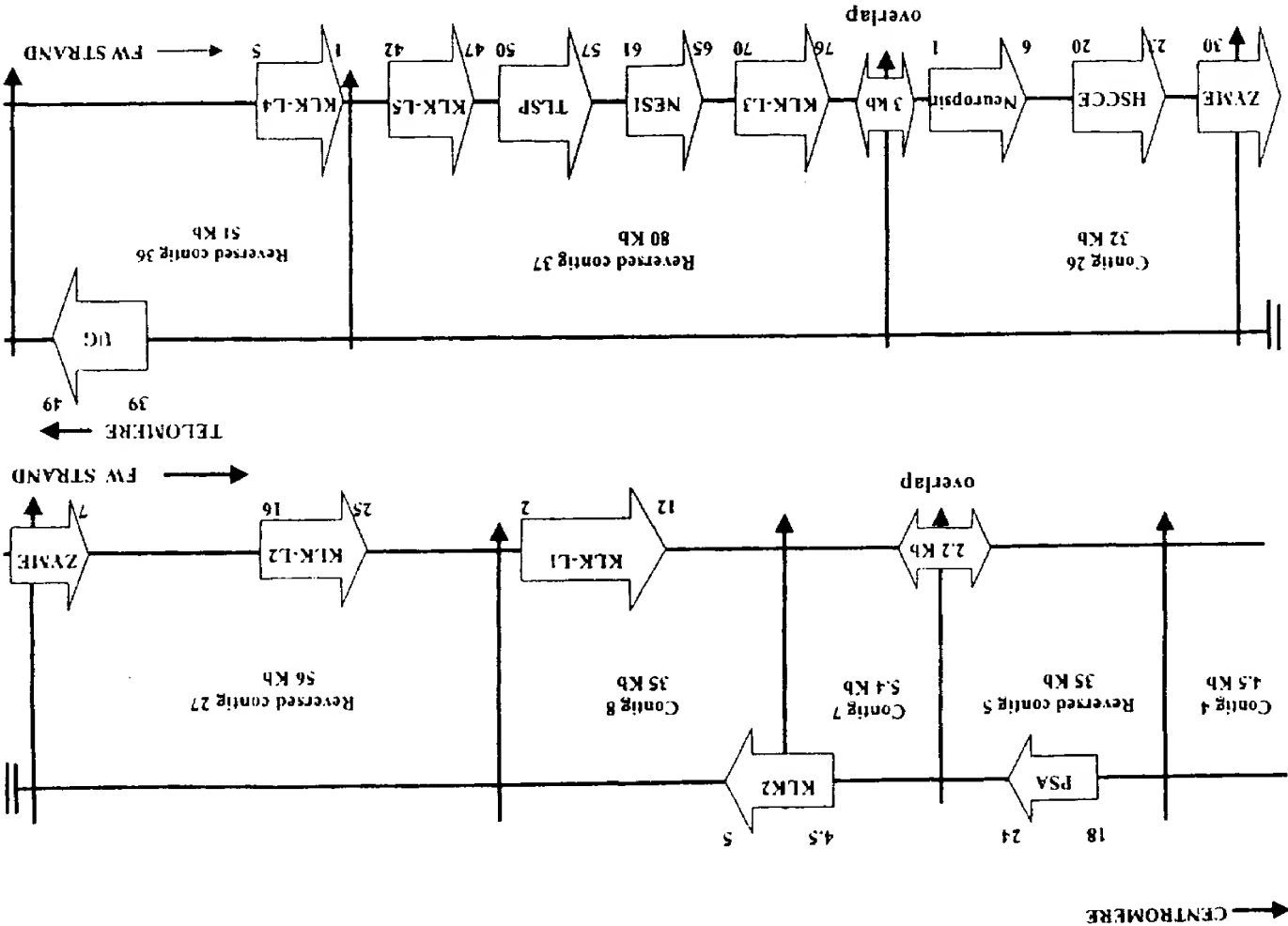
Identification of new genes. We have used a set of arbitrary rules to consider presence of a new gene in the genomic area of interest, as follows:

1. Clusters of at least 3 exons should be found.

(clones BAC 288H1 and BAC 76F7). These BACs were further analyzed by PCR and primers specific for PSA, NES1, KLK1 and KLK2. These analyses indicated that both BACs were positive for zymc, PSA and KLK2 and negative for KLK1 and NES1 genes.

Screening of the human PAC genomic library identified a PAC clone which was positive for NES1 (PAC 43B1). Further PCR analysis indicated that this PAC clone was positive for NES1 and KLK1 genes and negative for PSA, KLK2 and zymc. Combination of this information with the EcoRI restriction map of the region allowed us to establish the relative positions of these four genes. PSA is the most centromeric, followed by KLK2, zymc and NES1. Further

Figure 1. An approximate 300 Kb of contiguous genomic sequence around chromosome 19q13.3 - q13.4 represented by 8 contigs, each one shown with its length in Kb. The contig numbers refer to those reported in the Lawrence Livermore National Laboratory website. Note the localization of the seven known genes (PSA, KLK2, Zymc, NES1, HSCCE, neuropilin and TLSP) (see abbreviations for full names of these genes). All genes are represented with arrows denoting the direction of transcription. The gene with no homology to human kallikrein is termed UG (unknown gene). Numbers just below or just above the arrows indicate approximate Kb lengths in each contig. The length of each of these genes may change in the future since not all exons were identified for each new gene, as shown in Tables II-VII.



homologies with known human or animal kallikrein (D33). The five remaining genes all have significant (DB binding proteins 1 and 2 and with the surface antigen the kallikrein proteins. This gene has homology with the L(7) which showed no homology, at the protein level, with 1. In addition, we have identified one other new gene (gene protease (TLSP). Their relative location is shown in Figure the human neuropilin and the human trypsin-like serine human stratum corneum chymotryptic enzyme (HSCCE), analysis to be known genes not previously mapped, i.e. the genes of which three were found by subsequent homology

By using this strategy, we identified nine putative new prediction programs. they were identified by at least two different exon 3. We considered the exons predicted as reliable only if putative new genes. "excellent" quality, as indicated by the searching programs) were considered for the construction of the 2. Only exons with high prediction score ("good" or

Discussion

Prediction of protein-coding genes in newly sequenced DNA becomes very important after the establishment of

1. Conventional numbering of exons in comparison to the five coding exons of PSA, as described in Ref. 14.
2. Nucleotide numbers refer to the related contig (see text and Figure 1).
3. (+) = >95% homology with published human EST sequences.
4. Intron phase: 0 = the intron occurs between codons; 1 = the intron occurs after the first nucleotide of the codon; 2 = the intron occurs after the second nucleotide of the codon.
5. (+) denotes the exon containing the stop codon.
6. H = histidine, D = aspartic acid, S = serine. The amino acids of the catalytic triad are bold and underlined.
7. A = (neofinder (gene analysis), B = (neofinder (exon analysis), C = (trial 2), D = (GENEID-3).

Exon No. ¹	Putative coding region ²	From (bp)	To (bp)	No. of bases	Translated protein sequence	EST match ³	Intron phase ⁴	Stop codon ⁵	Catalytic triad ⁶	Exon prediction program ⁷
2	2263	2425	163	SLVSGSCSQINGEDCSPHSOP	WQVALVNIENETFCSGVLVH	+	II	-	II	A,B,D
3	2847	3109	263	NSYTGIGLHSLLEADQEPGSO	NIYVASTSVRIPEYVNPRLAND	+	I	-	D	A,B,C,D
4	3180	3312	137	GRMPTVYLQCVNVSVEEVS	KLYDPIYLHPSMFCAGGGQDDQ	+	0	-	-	A,B,C,D
5	4588	4737	150	GDSGGPLKNGYLGVSFGKA	PCGOVGFVVTNLCKFTTWIE	+	-	+	S	A,B,C

Table 11. Predicted exons of the putative gene KLK-L1. The translated protein sequences of each exon (open reading frame) are shown.

Table III. Predicted exons of the putative gene KLK-L2. The translated protein sequences of each exon (open reading frame) are shown*.

Exon No.	Putative coding sequence	No. of bases	Translated protein sequence	EST match ¹	Intron phase ⁴	Stop codon ⁵	Catalytic triad ⁶	Exon prediction ⁷	From (bp) To (bp)	
1	15,433	73	MATARPFWMMWVLCALITAI LGGT	+	1	-	-	-		
2	17,904	18,165	EHVIANNDVSCDIHPSNTVPSC SNODLGAGAGEDARSDSSSR INGSDCDMHTQPWQAVALLR PNOLYCGAVLVIFQWLLTAA HCRKK	+	II	-	H	A,B,C,D		
3	18,903	19,159	VFRVLGHVSLSPVYESGQMF QGVKSIPIHPGYSHPGHSNDML IKLNRRIRPTKDVRRPINVSSHCPS AGTKCI.VSGWGTTKSPQ	+	I	-	D	C,D		
4	19,245	19,378	VHPKVLQCLNISLSQKRCEDA YPRQIDDTMFCAAGDKAGRDSCQ	+	0	-	-	B,C		
5	24,232	24,384	GDSCGPPVNCNLSGLVSWGDY PCARPMPGVTNTLCKFTKWIOE TIOANS	+	-	+	S	A,B,C		

* All footnotes same as Table II.

Table IV. Predicted exons of the putative gene KLK-L1. The translated protein sequences of each exon (open reading frame) are shown*.

Exon No.	Putative coding region	No. of bases	Translated protein sequence	EST match ¹	Intron phase ⁴	Stop codon ⁵	Catalytic triad ⁶	Exon prediction ⁷	From (bp) To (bp)	
1	20,473	20,584	MEEEGDGMAYHKEALDA GCTQDP	-	I	-	-	A,B,C,D		
2	20,764	20,962	ACSSLPPLSLIPTPGHWAD TRAIAEECRPNQWQAG LFHLTRLCGATLISDRWLL TAAHCRK	+	II	-	H	A,B,C,D		
3	23,395	23,687	PLTSEACPSRYLWVRLGEHH LWKWEGPEQLFRVTDFEPHP GFNKDLSANDHNDIMLIRL PRQARLSPAVQPLNLSQTCVS PGMOCLISGWGAVSSPK	+	I	-	D	A,B,C,D		
4	26,305	26,441	ALFPVTLQCANISILNKICH WAYPGHISDSMLCAGLWEG GRGSCQ	+	0	-	-	A,B,C,D		
5	26,884	27,633	GDSCGPLVCNGLAGVSSGG AEPCSPRRPAAVYTSVCHYLD WIOEIMEN	-	-	+	S	A,B		

* All footnotes same as Table II.

large genome sequencing projects. This problem is the combination of potential functional signals with the global statistical properties of known protein-coding regions [15]. However, the most powerful approach for gene structure prediction is to combine information about potential functional signals (splice sites, translation start or stop signal etc.) together with the statistical properties of

* All footnotes same as Table II.

Exon No. ¹	Putative coding region ²	From (bp) To (bp)	No. of bases	Translated protein sequence	EST match ³	Intron phase ⁴	Stop codon ⁵	Catalytic triad ⁶	Exon prediction program ⁷
1	486	499	134	NPFDLLOCLNLSVSHATCIGV YPGRTSNMWCAGGVPQDQACQ	+	0	-	-	A,B,C,D
3	3592	3851	260	SRVWRLGEHSLSOLDWTEQ IRHSGISVTHPGYLGASTSHH DLRLRLRLPVRVTSSVOPPLP NDCATAGTECHVSGWGTNHPR	+	1	-	D	A,B,C,D
2	1588	1717	160	LSQVATPKFNNGTECGRNSQ PWQVGLFEGETSLRCGGVLID HRWVLTAHICSG	-	II	-	II	A,B,C

Table VI. Predicted exons of the putative gene K1.K-1.5. The translated protein sequences of each exon (open reading frame) are shown.*

* All footnotes same as Table II.

Exon No. ¹	Putative coding region ²	From (bp) To (bp)	No. of bases	Translated protein sequence	EST match ³	Intron phase ⁴	Stop codon ⁵	Catalytic triad ⁶	Exon prediction program ⁷
5	28,778	28,963	186	GDSCGPLVCNRLTYGIVSWGD FPCGPPDRPGVYTRVSRVYLV IRETRRKYEITOOQKWLRKGPQ	+	-	+	S	A,B,C
4	26,879	27,015	137	VNYPKTLQCANIQLRSDIECR OVPKGTIDNMLCAGTKKEGG KDSCE	+	0	-	-	A,B,C,D
3	25,460	25,728	269	GLKAYLGKHAIGRVEAGEQ YREVVHSHIPHEVRRSPHTN INDIDMHLJLQSPVQLTGVLQ TLP,SHNRLTPTCTRVSGW GTTTSPQ	+	1	-	D	A,B,C,D
2	24,915	25,120	176	ESSKVLNTNGTSGFLPGGYT CFPHSQPWQAALLVQGRLLC GGVLYHPKKWVLTAAHICLKE	+	II	-	II	C

Table V. Predicted exons of the putative gene K1.K-1.4. The translated protein sequences of each exon (open reading frame) are shown.*

coding sequences (coding potential) along with information about homologies between the predicted protein and already known protein families [16]. In mouse and rat, kallikreins are encoded by large multigene families and these genes tend to cluster in groups with a distance as small as 3.3 - 7.0 Kb [3]. A strong conservation of gene order between human chromosome 19q13.1 - q13.4 and 17 loci in a 20-cM proximal part of mouse chromosome 7, including the kallikrein locus, has been documented [17].

In humans, only a few kallikrein genes were identified. In fact, only KLK1, KLK2 and KLK3 (PSA) are considered to represent the human kallikrein gene family [9,18]. In this paper, we provide strong evidence that a large number of kallikrein-like genes are clustered within a 300Kb region around chromosome 19q13.2 - q13.4. The three established human kallikreins (KLK1, KLK2, KLK3), zyme and NES1, around chromosome 19q13.2 - q13.4. The three established human kallikreins (KLK1, KLK2, KLK3), zyme and NES1, constitute a family that likely originated by duplication of an ancestral gene. The relative localization of all these

as well as the stratum corneum chymotryptic enzyme, neuropsin and TLSF (trypsin-like serine protease) and another five new genes, KLK-L1 to KLK-L5, may constitute a large gene family. This will bring the total number of kallikrein or kallikrein-like genes in humans, in this region of chromosome 19, to thirteen.

The human stratum corneum chymotryptic enzyme [19], are three previously characterized genes which have many structural similarities with the kallikreins and other members of the serine protease family. However, they have not been mapped in the past. Our precise mapping of all three genes in the region of the kallikrein gene family further suggests that these genes, along with the ones that were newly identified by us, and the already known ones, constitute a family that likely originated by duplication of an ancestral gene. The relative localization of all these

* All footnotes same as Table II.

Exon No.	Putative coding region ¹	No. of bases	Translated protein sequence	EST match ²	Intron phase ³	Stop codon ⁴	Catalytic program ⁵
1	44,129	44,641	513	+	1	-	B.C.
			PPLSLFPAVPERRTLRNRRSLAALAPL				
			TPDMLLLPLWGRRERAEQTSKLL				
			TMOSSTVQEGLCVHVPCSFSPSHG				
			WYFPGPVVHGWFREGANPDQAPV				
			ATNNPAAVWEETRDRETHLGDPTTK				
			NCTLSIRDAKRSADAGRYFFERKIEKGSIK				
			WNYKHHRLSNVNT				
2	44,813	45,121	279	+	1	-	A,B,C,D
			ALTHRPNIILPGTLFSGCPQNLTCVPPW				
			ACEQGTTPPMISWIGTSVSPLDPSTTRSSV				
			LTLIPQPDHGTSLTCQVTFPGASVTTN				
			KTVHLNVS				
3	45,327	45,374	48	-	1	-	A,B,D
			YPPQNLTMVTFQGDGT				
4	46,318	46,542	225	+	1	-	A,B,C
			EGOSRLRLVCADVADSNPPARLSLWKR				
			GLTLCPSQSPNPGVLELPWHLRDAAE				
			FTCRANPLGSOQVYLVNSLO				
5	47,195	47,283	186	+	0	-	A,B,C,D
			SKATSGVTQGVVGAGATATLFLSFC				
			VIFV				
6	49,136	49,554	186	+	-	+	A,B,C
			GPLTFPWAEDSPDPPPASARSSVGE				
			GELQYASLSFQMKPWS RGQEATD				
			TEYSSEIKHHR				

Table VII. Predicted exons of the unknown gene UCI. The translated protein sequences of each exon (open reading frame) are shown.*

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Table VIII. Homology between the predicted amino acid sequences of the newly identified putative genes and protein sequences deposited in EMBL.

No. Cene	Homologous known protein	Identity (%)	amino acids
1	KLK-L1	• Human stratum corneum chymotryptic enzyme 44 (101/227)	
2	KLK-L2	• Human neuropilin 48 (106/219) • Human stratum corneum chymotryptic enzyme 47 (103/210)	
3	KLK-L3	• Human procathepsin M 45 (99/219) • Human trypsinogen 1 45 (100/221) • Rat trypsinogen 44 (98/220)	
4	KLK-L4	• Human procathepsin M 52 (118/225) • Human neuropilin 51 (116/225) • Mouse neuropilin 51 (116/226) • Human glandular kallikrein 48 (113/234) • Human prostatic specific antigen 47 (108/227)	
5	KLK-L5	• Human neuropilin 44 (81/184) • Rat trypsinogen 1 42 (76/178) • Rat trypsinogen II 42 (75/178) • Human procathepsin M 41 (73/178)	
6	UG	• Human myeloid cell surface antigen CD33 61 (144/233) • Human OB binding protein-2 50 (166/328) • Human OB binding protein-1 43 (189/431) • Human myelin associated glycoprotein 27 (86/311)	

genes is depicted in Figure 1. We consider Figure 1 to describe the human kallikrein gene family, consisting of

Kallikrein genes are a subfamily of serine proteases, thirteen genes.

bradykinin (kallidin) from kininogen [22]. More recently, however, a new, structural concept has emerged to describe kallikreins. From accumulated sequence data, it is now clear that the mouse has many genes with high homology to kallikrein coding sequences [23-24]. Richards and co-workers have contributed to the concept of a "kallikrein multigene family" to refer to these genes [25-26]. This definition is not based much on specific enzymatic function of the gene product, but more on its sequence homology and their close linkage on mouse chromosome 7. In humans, only KLK1 meets the functional definition of a kallikrein. KLK2 has trypsin-like enzymatic activity and KLK3 (PSA) has very weak chymotrypsin-like enzymatic activity. These activities of KLK2 and KLK3 are not known to liberate biologically active peptides from precursors. Based on the newer definition, members of the kallikrein family include, not only the gene for the kallikrein enzyme, but also genes encoding other homologous proteases, including the enzyme that processes the precursors of the nerve growth factor and epidermal growth factor [8]. Therefore, it is important to note the clear distinction between the enzyme kallikrein and a kallikrein or a kallikrein-like gene.

It is important to mention that the prediction of new

genes by computer programs is still not a straightforward process. Many shortcomings are known to exist in such programs. Most of these programs are unable to detect non-coding exons and non-coding portions of exons. Some programs are unable to detect exons without the presence of a genomic context (when the regions adjacent to an exon are not present). Also, the power for detection of small exons (less than 100 bp) is low in some programs. About 5% of real splice sites are usually lost by some programs but over-prediction is usually small [27]. However, the detection power of some programs (e.g. Gital 2) is about 91% when tested with known genomic sequences. An indication of the quality of prediction is provided with these programs. In our study, we considered only exons which were predicted with "good" or "excellent" quality and only exons which were predicted by at least two different programs. Moreover, we considered the presence of a putative gene only when at least three exons clustered coordinately in that region. Additional evidence that these new genes are indeed homologous to the known kallikreins and other serine proteases comes from comparison of the intron phases. As we have published previously [14], trypsinogen, PSA and NES1 have 5 coding exons of which the first has intron phase I (the intron occurs after the first nucleotide of the codon), the second has intron phase II (the intron occurs after the second nucleotide of the

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codon), the third has intron phase I and the fourth has fifth exon contains the stop codon. The intron phases of the predicted new kallikrein-like genes follow these rules and are shown in the respective tables. Further support comes from our identification in the new genes, of the conserved amino acids of the catalytic domain of the serine proteases, as presented in Tables II - VI.

In order to test the accuracy of the gene prediction programs, we tested known genomic areas containing the PSA, zyme and KLK2 genes. Two of these programs (Grail 2 and GeneBuilder) were able to detect about 95% of the tested known genes (data not shown). Matches with expressed sequence tag sequences (EST) can also be employed for gene structure prediction in the GeneBuilder program and this can significantly improve the power of the program, especially at high stringency (e.g. >95% homology). In the respective Tables for each putative new gene we provide evidence for matching ESTs from the Genbank human EST database. The presence of EST matches is additional strong evidence that these newly identified genes are expressed.

The question remains if these new genes are functional. In mouse, ten of the kallikrein genes appear to be pseudogenes [9]. One of our new genes (UG) does not show homology with the kallikrein genes. However, it has some protein homology with myelin associated glycoprotein (Table VIII). There may still be an association between UG and the kallikrein genes since some mouse kallikreins are related to nerve growth factor, as discussed earlier [8] and zyme as well as neuropilin and TLSP were found to be highly expressed in brain tissue and is claimed that zyme may be related to Alzheimer's disease [11]. We are now screening and sequencing EST clones and studying the tissue expression of these new genes by RT-PCR. Our goal is to fully characterize their mRNA sequence, study their expression and regulation and examine if they are involved, or can be used, like other human kallikreins (e.g. PSA, KLK2, zyme and NES1), in breast, prostate, testicular or other cancer diagnostic, prognostic or therapeutic applications. The expansion of the kallikrein locus in humans to thirteen genes will allow us to better understand the role of this family in various cancers. There is already evidence that some of these genes encode for tumor suppressors [10, 28, 29].

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